## Philippe D'Arco

List of Publications by Year in descending order

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		933447	839539
18	1,550	10	18
papers	citations	h-index	g-index
10	1.0	10	2270
18	18	18	2370
all docs	docs citations	times ranked	citing authors

#	Article	IF	CITATIONS
1	The superexchange mechanism in crystalline compounds. The case of KMF <sub>3</sub> (M = Mn, Fe, Co,) Tj ETQo	q1.1 0.784 1.8	4314 rgBT  €
2	Strategies for the optimization of the structure of crystalline compounds. Journal of Computational Chemistry, 2022, 43, 184-196.	3.3	9
3	Interstitial carbon defects in silicon. A quantum mechanical characterization through the infrared and Raman spectra. Journal of Computational Chemistry, 2021, 42, 806-817.	3.3	2
4	The ferromagnetic and anti-ferromagnetic phases (cubic, tetragonal, orthorhombic) of KMnF <sub>3</sub> . A quantum mechanical investigation. Physical Chemistry Chemical Physics, 2021, 23, 26780-26792.	2.8	7
5	From anisotropy of dielectric tensors to birefringence: a quantum mechanics approach. Rendiconti Lincei, 2020, 31, 835-851.	2.2	3
6	The CRYSTAL code, 1976–2020 and beyond, a long story. Journal of Chemical Physics, 2020, 152, 204111.	3.0	133
7	Substitutional carbon defects in silicon: A quantum mechanical characterization through the infrared and Raman spectra. Journal of Computational Chemistry, 2020, 41, 1638-1644.	3.3	8
8	Vibrational spectroscopy of hydrogens in diamond: a quantum mechanical treatment. Physical Chemistry Chemical Physics, 2018, 20, 11930-11940.	2.8	17
9	The Infrared spectrum of very large (periodic) systems: global versus fragment strategiesâ€"the case of three defects in diamond. Theoretical Chemistry Accounts, 2018, 137, 1.	1.4	10
10	Hydrogen, boron and nitrogen atoms in diamond: a quantum mechanical vibrational analysis. Theoretical Chemistry Accounts, 2018, 137, 1.	1.4	16
11	On the use of the symmetry-adapted Monte Carlo for an effective sampling of large configuration spaces. The test cases of calcite structured carbonates and melilites. Computational Materials Science, 2017, 126, 217-227.	3.0	2
12	Hydrogrossular, Ca <sub>3</sub> Al <sub>2</sub> (SiO <sub>4</sub> ) <sub>3–x</sub> (H <sub>4</sub> O <sub>4</sub> ) <sub>x<an 100,="" 2015,="" 2637-2649.<="" ab="" american="" and="" energetic="" initio="" investigation="" its="" mineralogist,="" of="" properties.="" structural="" td=""><td></td></an></sub> :		16
13	C <scp>RYSTAL14</scp> : A program for the <i>ab initio</i> investigation of crystalline solids. International Journal of Quantum Chemistry, 2014, 114, 1287-1317.	2.0	1,151
14	On the use of symmetry in configurational analysis for the simulation of disordered solids. Journal of Physics Condensed Matter, 2013, 25, 105401.	1.8	34
15	Symmetry and random sampling of symmetry independent configurations for the simulation of disordered solids. Journal of Physics Condensed Matter, 2013, 25, 355401.	1.8	24
16	On the use of symmetry in the <i>ab initio</i> quantum mechanical simulation of nanotubes and related materials. Journal of Computational Chemistry, 2010, 31, 855-862.	3.3	48
17	Structure and energetics of imogolite: a quantum mechanical ab initio study with B3LYP hybrid functional. Journal of Materials Chemistry, 2010, 20, 10417.	6.7	41
18	Single-layered chrysotile nanotubes: A quantum mechanical <i>ab initio</i> simulation. Journal of Chemical Physics, 2009, 131, 204701.	3.0	26